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71) Applicant: Montedison S.p.A. 31, Foro Buonaparte 1-20121 Milan(IT)

(72) Inventor: Parodi, Sandro 4, via dei Negri Oleggio Novara(IT)

(72) Inventor: Nocci, Roberto 170, corso XXIII Marzo Novara(IT)

(72) Inventor: Giannini, Umberto, Dr. 53, via Sismondi Milano(IT)

(72) Inventor: Barbe, Pier Camillo, Dr. 3, via Pavone Ferrara(IT)

(72) Inventor. Scata, Umberto 31, via C. Battisti Ferrara(IT)

(14) Representative: Zumstein, Fritz, Dr. Dr. F. Zumstein sen. Dr. E. Assmann et al, Dr. R. Koenigsberger Dr. F. Zumstein jun. Dipl.-Ing. F. Klingseisen Brauhausstrasse 4 D-8000 München 2(DE)

(54) Components and catalysts for the polymerization of olefins.

(57) Catalysts for the polymerization of alpha-olefins which comprise the reaction product of:

a) an Al-aikyl compound;

b) a silicon compound containing at least one Si-OR or Si-OCOR or Si-NR₂ bond, R being a hydrocarbyl radical;

c) a solid comprising as essential support, a Mg dihalide in active form and supported therein a Ti halide or a halo-Ti-alcoholate and a particular type of electron donor compound.

DESCRIPTION

The present invention refers to new supported components of catalysts for the polymerization of CH₂=CHR olefins wherein R is an alkyl radical with 1 to 4 carbon atoms, or an aryl radical, and mixtures of said olefins with ethylene and the catalysts obtained from said components.

The supported highly active and highly stereospecific catalysts up to now known for the polymerization of propylene and higher olefins, are obtained by the reaction of an Al-alkyl compound partially complexed with an electron donor compound (outside donor) with a solid component comprising a Ti compound and an electron-donor compound (inside donor) supported on a Mg-halide in active form.

Examples of such catalysts have been described in British Patent No. 1,559,194 and Belgian Patent No. 868,682.

Outside donor consisting of silicon compounds containing Si-O-C bonds have been also described (Japanese patent application Sho 79/94590 and Sho 80/36203).

Among the various and numerous inside donors also com-

pounds as methyl methacrylate and ethyl pivalate have been cited.

However in all the prior art catalysts in which a silicon compound containing Si-O-C bonds is used as outside donor, esters of benzoic acid and derivates thereof are used as inside donor.

The performance of the above catalysts, expressed in terms of activity and stereospecificity is not different from the performance of the catalysts in which ethyl ben zoate and similar esters of benzoic acid are used as outside donor.

It has now unexpectedly been found that it is possible to increase the activity and stereospecificity of the prior art supported catalysts comprising as outside donor a silicon compound containing Si-O-C bonds by using as inside donor an ester having a particular structure.

The catalysts of this invention comprise the product of reaction between the following components:

an Al-trialkyl or an Al-alkyl compound containing

2 or more aluminum atoms linked to each other

through oxygen or nitrogen atoms or through SO₄ or

SO₃ groups;

- b) a silicon compound containing one or more Si-OR, Si-OCOR or Si-NR₂ bonds (R being a hydrocarbyl radical);
- c) a solid comprising, as essential support, an anhydrous Mg-dihalide present in active form, and supported on said dihalide a Ti-halide or a Ti-haloalcoholate and an electron-donor compound selected from the following groups of compounds:
 - 1) Mono and polyesters of saturated polycarboxylic acids wherein at least one of the esteric carbonyl groups is linked to a tertiary or quaternary carbon atom or to a linear or branched chain of at least 4 carbon atoms;
 - 2) Mono and polyesters of unsaturated polycarboxylic acids wherein two carboxy groups are linked to vicinal double bond-forming carbon atoms and in which at least one of the R hydrocarbyl radicals of the COOR groups is a branched saturated or unsaturated radical with 3 to 20 C atoms or is an aryl or arylalkyl radical with 6 to 20 C atoms;
 - 3) Mono and diesters of aromatic bicarboxylic acids having the COOH groups in ortho position wherein

- at least one of the R hydrocarbyl radical of the COOR groups contains from 3 to 20 carbon atoms;
- 4) Mono and polyesters of aromatic hydroxy compounds containing at least 2 hydroxyl groups in ortho position;
- a hydroxyl group is in ortho position to the carboxy group;
- acids wherein at least one of the hydrocarbyl R and R' radicals of the R COOR' group is a saturated or unsaturated branched radical containing from 3 to 20 C atoms, or is an arylalkyl radical with 7 to 20 C atoms or R is an aryl radical with 3 to 20 carbon atoms linked to the esteric carbonyl group directly or through a methylene group. and in which the R' radical contains from 3 to 20 C when it is a linear hydrocarbyl radical;
- 7) Esters of carbonic acid of formula CO in OR which at least one of the R radicals which can be the same or different are hydrocarbyl radicals with 3 to 20 carbon atoms.

Representative esters which are suitable in preparing component c) are the following:

Class 1)

diethyl diisobutylmalonate, diethyl n-butylmalonate, diethyl-n-dibutylmalonate, diethylphenylmalonate, diethyl-1,2--cyclohexane-dicarboxylate, dioctylsebacate, diisobutyl adipate.

Class 2)

di-2-ethyl-hexyl-maleate, diisobutylmaleate, diisobutyl-3,4-furan-dicarboxylate, di-2-ethylhexylfumarate, 2-ethylhexylmonomaleate.

Class 3)

diisobutyl-2,3-naphthalen-dicarboxylate, di-n-propyl, di-n-butyl, diisobutyl, di-n-heptyl, di-2-ethyl-hexyl, di-n-octyl, di-neopentil phthalates, monobutyl and monoisobutyl esters of phthalic acid, ethyl-isobutyl-phthalate, ethyl-n-butyl-phthalate.

Class 4)

2,3-diacetoxynaphthalene, 1,2-diacetoxybenzene, 1-methyl--2,3-diacetoxybenzene.

Class 5)

Benzoyl-ethylsalicylate, acetyl-methylsalicylate.

Class 6)

Ethylene-glycol-pivalate, 1,4-butanediol-pivalate, benzyl and isobutylpivalate, n-propylpivalate, ethyl diphenyl-acetate, isobutylmethacrylate, isobutylacrylate, ethyl-benzoilacetate, isobutylpyruvate, isobutyl-trans-3-methoxy-2-butenoate.

Class 7)

phenyl-ethylcarbonate, diphenyl carbonate.

Preferred compounds are the esters of maleic, pivalic, methacrylic, carbonic and phthalic acids.

As indicated, the esters of the polycarboxylic acids can contain beside the ester groups also unesterified COOH groups.

In preparing component c) the esters are contacted with the active Mg dihalide or the precursors of said dihalides as preformed compounds or the esters can be formed in situ by means of known reactions as for instance by esterification between an alcohol or an alcoholate and an aryl halide or between an anhydride or an emiester of a polycarboxylic acid with an alcohol or by transesterification. The esters can be also used in mixture with other known inside donors.

The active anhydrous Mg dihalides forming the essential support of component c) are the Mg dihalides showing in

the X-ray powder spectrum of component c) a broadening of at least 30% of the most intense diffraction line which appears in the powder spectrum of the corresponding dihalide having 1 m^2 /g of surface area or are the Mg dihalides showing a X-ray powder spectrum in which said most intense diffraction line is replaced by a halo with the intensity peak shifted with respect to the interplanar distance of the most intense line and/or are the Mg dihalides having a surface area greater than 3 m^2/g .

The measurement of the surface area of the Mg dihalides is made on component c) after treatment with boiling TiCl4 for 2 hours. The found value is considered as surface area of the Mg dihalide.

Very active forms of Mg dihalides are those showing a X-ray powder spectrum in which the most intense diffraction line appearing in the spectrum of the corresponding halide having 1 m²/g of surface area is decreased in relative intensity and broadened to form a halo or are those in which said most intense line is replaced by a halo having its intensity peak shifted with respect to the interplanar distance of the most intense line. Generally the surface area of the above forms is higher than 30-40 m²/g

and is comprised in particular between $100-300 \text{ m}^2/\text{g}$.

Active forms are also those deriving from the above forms by heat-treatment of component c) in inert hydrocarbon solvents and showing in the X-ray spectrum sharp diffraction lines in place of the halos.

The sharp, most intense line of these forms shows in any case a broadening of at least 30% with respect to the corresponding line of the Mg dihalide having 1 m²/g of surface area. Preferred Mg dihalides are Mg dichloride and Mg dibromide. The content in water of the dihalides is generally less than 1% by weight.

For Ti halides or Ti haloalcoholates and esters supported on active Mg dihalide is meant the above compounds which may be chemically or physically fixed on the support, not extractable from component c) by treatment of the same with boiling 1,2-dichloroethane for 2 hours.

Components a), b) and c) are made to react each other in any order; preferably, however, components a) and b) are premixed before being contacted with component c).

Component c) may be premixed with either component

a) and/or b). The pre-mixing of a) and b) is conducted at temperatures comprised, usually, between room
temperature and the temperature used in the polymerization process.

The pre-reaction of c) and b) may be carried out also at higher temperatures. Compound b) may be also in-corporated and made to react with component c) itself.

Component b) is made to react in a molar ratio with respect to the halogenated Ti compound supported on component c) of at least 1 and in a molar ratio with respect to the Al-alkyl compound used as component a) of less than 20 and preferably comprised between 0.05 to 0.3.

In component c) the molar ratio between the Mg dihalide and the halogenated Ti compound supported therein is comprised between 1 and 500 and the molar ratio between said halogenated Ti compound and the electron-donor supported on the Mg dihalide is comprised between 0.1 and 50. The silicon compounds set forth in b) include compounds of general formula:

R SiY X p

wherein:

- R is an alkyl, alkenyl, aryl, arylalkyl, cycloalkyl radical with from 1 to 20 carbon atoms;
- Y is a -OR', -OCOR', -NR2' wherein R', either equal to or different from R, has the same meaning as R;
- X is either a halogen or hydrogen atom or a -OCOR", $-NR_2$ " group wherein R", either equal to or different from R', has the same meaning as R';

m, n and p are numbers comprised respectively between:

m between 0 and 3, n between 1 and 4 and p between 0 and
1; and m+n+p is equal to 4.

Other silicon compounds that may be used are compounds in which two or more silicon atoms are bound to each other through oxygen or nitrogen atoms.

Examples of these compounds are hexaethoxydisiloxane, simmetrical diphenyltetraethoxydisiloxane

$$(c_2^{H_5^{O}})_2^{-\text{Si-O-CH}}_2^{-\text{CH}}_2^{-\text{O-Si-(OC}}_2^{H_5})_2$$
 $c_6^{H_5}$

Preferred silicon compounds are: phenylalkoxysilanes as phenyltriethoxy or trimethoxysilane, diphenyldimethoxy and diethoxysilane, monochlorophenyldiethoxysilane; alkylalkoxysilanes as ethyltriethoxysilane, ethyltriisopropoxysilane.

Examples of other suitable compounds are: chlorotrie-

thoxysilane, acetoxytriethoxysilane, vinyltriethoxysilane, butyltriethoxysilane, triphenylmonoethoxysilane, phenyltricycloethoxysilane, phenyldiethoxydiethylaminosilane, tetraphenoxysilane or tetralkoxysilanes as tetramethoxysilane.

The silicon compound can be also formed in situ by reaction for instance of a halogenated silicon compound as SiCl_4 with an alcohol or an alcoholate of Mg or Al.

In the catalysts of the invention the silicon compound is present in a combined form in the solid product
of the reaction between the various catalyst- forming components, in a molar ratio between the silicon compound and
the halogenated Ti compound greater than 0.05 and generally
comprised between 0.1 and 5.

The Al-alkyl compounds forming component a) includes Al-trialkyls as for instance Al-triethyl, Al-triisobutyl, Al-triisopropyl and compounds containing two or more Al atoms linked to each other through etero-atoms as:

$$(c_{2}^{H_{5}})_{2}^{\Lambda 1-O-\Lambda 1}(c_{2}^{H_{5}})_{2}^{2}$$
 $\cdots (c_{2}^{H_{5}})_{2}^{\Lambda 1-N-\Lambda 1}(c_{2}^{H_{5}})_{2}^{2}$
 $(c_{2}^{H_{5}})_{2}^{\Lambda 1-O-S-O-\Lambda 1}(c_{2}^{H_{5}})_{2}^{2}$

As indicated λ 1-alkyl compounds in which λ 1 atoms are linked through groups as SO₄ or SO₃ are also suitable.

The Al-alkyl compounds may be used in mixture with Al-alkyl halides as AlEt₂Cl.

Component c) is prepared according to known methods.

One of these methods consists in co-milling the Mg halide and the electron-donor compound of this invention until the appearance in the X-ray spectrum of the milled product of the modifications above set forth for the spectrum of the Mg dihalide and thereafter reacting the milled product with the Ti-compound.

Preparations of this type are described in British Patent No. 1,559,194.

Similar preparations are described in U.S. Patents Nos. 4,107,413, 4,107,414 and 4,107,415.

Another method consists in reacting the adduct of a Mg halide with an alcohol, with a Ti compound in the presence of an electron-donor compound not containing active hydrogen atoms. This method is described in Belgian Patent No. 868,682.

According to another method, which is described in published German application 3,022,738, the adduct between the Mg dihalide and the alcohol is reacted in liquid

form with the halogenated Ti compound and the electron-donor compound.

Further methods are described in published German application 2,924,029, U.S. Patent 4,220,554 as well as in Italian Patent Ap plication No. 27.261/79.

Another method consists in co-milling the Mg dihalide, the halogenated Ti compound and the electron-donor compound until activation of the Mg dihalide and in treating a suspension of the milled product in a halogenated hydrocarbon as 1,2-dichloroethane, chlorobenzene, methylene chloride, hexachloroethane.

The treatment is carried out at temperatures comprised between 40°C and the boiling point of the halogenated hydrocarbon for a time ranging in general from 1 to 4 hours.

According to another method a porous support like $5iO_2$ or Al_2O_3 , having a low content of OH groups (preferably less than 1% by weight) is impregnated with a liquid adduct between the Mg dihalide and an alcohol; the support is then treated with an excess of $TiCl_4$ containing dissolved the electron-donor compound according to the procedure described for instance in German patent application 3,022,728 or Belgian patent 868,682.

In all the above methods the final product contains a Mg dihalide, present in the active form as set forth here above.

Other known methods which lead to the formation of Mg dihalide in active form or to Ti- containing Mg dihalide supported components, in which the dihalide is present in active form, are based on the following reactions:

- reaction of a Grignard reagent or a Mg R₂ compound

 (R being a hydrocarbyl radical) or complexes of

 said Mg R₂ compounds with Al-trialkyls, with halogenating agents as AlX₃ or Al R_M compounds (X is halogen, R is a hydrocarbyl m+n = 3), SiCl₄ or HSiCl₃;
- reaction of a Grignard reagent with a silanol or polysiloxane, H₂O or with an alcohol and further reaction with a halogenating agent or with TiCl₄;
- reaction of Mg with an alcohol and a halogenidric acid or of Mg with a hydrocarbyl halide and an alcohol;
- reaction of MgO with Cl₂ or AlCl₃;
- reaction of MgX2.nH2O (X = halogen) with a halogenating agent or TiCl4;

reaction of Mg mono or dialcoholates or Mg carboxylates with a halogenating agent.

The Ti-halides or Ti-halogenalcoholates, include in particular the Ti-tetrahalides, Ti-trihalides and Ti-trihalogenalcoholates. Preferred compounds are: TiCl₄, TiBr₄, 2,6-dimethylphenoxytrichlorotitanium.

methods, for instance by reduction of TiCl₄ with Al or a metallorganic Al compound or with hydrogen.

In case of the Ti-trihalides it may be convenient for the purpose of improving the performance of the catalysts to carry out an oxidization, even if partial, of the titanium, either during or after the preparation of component c).

To this purpose there may be used halogens, iodine halides.

Preferred catalysts are those in which: component c) is obtained from MgCl₂, TiCl₄ and esters of maleic, pivalic and phthalic acids and in which component b) is phenyl or ethyltriethoxysilane or diphenyldimethoxy or diethoxysilane.

Component a) is an Al-trialkyl as Al-triethyl or Al-triisobutyl.

Component c) is prepared according the methods described in Brit. Pat. No. 1,559,194, Belgian patent No. 868,682, published German application No. 2,924,029, U.S. Patent 4,220,554, Italian Pat. Appln. 27,261/79 or published German application 3,022,738.

The preferred method of preparing component c) includes also the co-milling of MgCl₂, TiCl₄ and the ester and in treating the milled product with a halogenated hydrocarbon as 1,2-dichloroethane.

The catalysts according to the invention are used to polymerize the alpha-olefins according to known methods that is, in carrying out the polymerization in a liquid phase, either in the presence or absence of an inert hydrocarbon solvent, or in gas phase or also by combining, for instance, a liquid phase polymerization step with a step in gas phase.

In general the temperature is comprised between 40° and 160°C, but preferably between 60° and 90°C, operating either at atmospheric or at greater than atmospheric pressure.

As a molecular weight regulator hydrogen or other regulators of a known type are used.

The catalysts are used particularly suitable in polymerizing propylene, butene-1, styrene, 4-methylpentene. The catalysts may also be used according known methods to polymerize mixtures of propylene and ethylene to form modified polypropylenes having better shock-resistance at low temperatures (the so called block copolymers of propylene and ethylene) or to obtain random crystalline copolymers of propylene with minor proportions of ethylene.

The following examples are given for merely illustrative purpose and are not intended to be in any way limiting the scope of the invention.

EXAMPLES 1 - 10

Into a stainless steel autoclave having a total volume of 3cl, equipped with a magnetical stirrer and a thermocouple heat-stabilized at 60°C and kept under pressure by a nitrogen atmosphere, there were introduced 1000 ml of a suspension in degassed and anhydrous n-heptane containing 5 m mols of triethylaluminum, the phenyltriethoxysilane (PES)

and a solid catalytic component prepared according to example 1 of Italian Pat. Appln. 27.261/79, but using, instead of ethylbenzoate, the esters listed in Table I, while propylene was also fed in. The catalytic components thus prepared showed X-ray powder spectra in which the most intense diffraction line appearing in the spectrum of MgCl₂ having 1 m²/g of surface area is decreased in relative intensity and broadened to form a halo.

The autoclave was closed and thereafter hydrogen was introduced up to a pressure of 0,2 atmospheres, the temperature was brought to 70°C and simultaneously, propylene was introduced up to a total pressure of 7 atmospheres.

During the polymerization the pressure was kept constant by continuous feeding of the monomer. After 4 hours, the polymerization was stopped by quick cooling and degassing of the polymeric slurry. The polymer was separated from the solvent by filtering and was dried in a hot nitrogen flow at 70°C. The quantity of polymer dissolved in the filtrate was thereupon isolated, weighed and summed to the polymer soluble in boiling n-heptane, for the calculation of the isotacticity index (I.I.).

The quantity of catalytic component used and the content of Ti in said component, the molar ratio of the phenyltrietoxysilane with respect to the triethylaluminium, the yield in polymer with respect to the introduced catalytic component, the isotacticity index (I.I.), the surface area of the solid catalytic component and the inherent viscosity determined in tetralin at 135°C, are reported in Table I.

EXAMPLE 11

Example 8 was repeated under the same polymerization conditions described in Examples 1-10, but using a solid catalytic components prepared according to Example 7 of German Pat. Appln. 2.643.143.

The catalytic component showed a X-ray powder spectrum in which the most intense diffraction line appearing in the spectrum of MgCl₂ having 1 m²/g of surface area is decreased in relative intensity and broadened to form a halo.

The content of Ti in the solid catalytic component, the quantity of said component, the molar ratio between triethylaluminium and PES, the results of the polymerization test are reported in Table I.

EXAMPLE 12

The solid catalytic component prepared according to Exam-

ple 11 was suspended in the inert hydrocarbon solvent

Isopar G (a mixture of isoparaffinic hydrocarbon boiling
in the temperature range of from 158° to 172.5°C) and
heat-treated at 120°C for 24 hours. The catalytic components thus obtained showed a X-ray powder spectrum in
which the most intense diffraction line appeared as a
sharp line, the half peak breadth of which however was
broadened more than 30% the corresponding half peak breadth
of MgCl₂ having 1 m²/g of surface area.

The catalyst component obtained was used under the same polymerization conditions described in example 11.

The characteristics of the solid catalytic component, the results of the polymerization test are reported in Table I.

Comparative EXAMPLES 1 and 2

Example 1 was repeated but using the solid catalytic component prepared according to example 1 of Italian Pat. Appl.

No. 27,261/79. The X-ray powder spectrum was similar to that of the catalytic component of Example 1.

The results of the polymerization tests have been reported in Table I.

Comparative EXAMPLE 3

Example 11 was repeated except that the solid catalytic component prepared according to Example 7 of German Pat.

Appln. 2,643,143.was used. The X-ray powder spectrum was similar to that of catalytic component of Example 11.

The results of the polymerization tests have been reported in Table I.

EXAMPLES 13-18

Example 1 was repeated using a solid catalytic component prepared as follows.

Anhydrous MgCl₂, an ester, listed in Table II, and TiCl₄ in molar ratio of 1:1 with respect to the ester, were co-ground in a vibrating mill of the type VIBRATOM manufactured by N.V. TEMA'S, Gravenhage (Holland), having a total volume of one liter and containing 3 kg of stainless steel balls of 16 mm diameter.

Grinding was effected for 72 hours employing a filling $coe\underline{f}$ ficient equal to 100 g/l of total volume (vacuum), at an interior temperature of the mill of $25^{\circ}C$.

Charging of the mill, the grinding and discharging of the mill occurred in a nitrogen atmosphere.

10 g of the co-ground product was contacted with 100 ml of 1,2-dichlorcethane at 80° C for 2 hours.

After this period, 1,2-dichloroethane was removed by filo tration at 80 C and the residual solid product was repeatedly washed with n-heptane at room temperature till the disappearance of the chlorine ions from the filtrate and then was kept in heptanic suspension.

The catalytic components thus prepared showed a X-ray powder spectrum in which the most intense diffraction line appearing in the spectrum of MgCl₂ having 1 m^2/g of surface area was decreased in relative intensity and broadened to form a halo.

The type of ester, the characteristics of the solid catalytic component, and the results of polymerization tests have been reported in Table II.

EXAMPLES 19 - 26

Example 1 was repeated but using a solid catalytic component prepared according to Example 3 of Italian Patent Appln. No. 26,908 A/78.

A solid adduct MgCl $_2.2.5$ C $_2$ H $_5$ OH in the form of spherical particles was slowly added to a suspension of an adduct TiCl $_4$ -ester using molar ratios Mg/ester of 10 and TiCl $_4$ /C $_2$ H $_5$ OH of 10.

The whole was then heated at 100°C, kept at said temperature for 2 hours and then filtered at 100°C. The resulting solid product was treated with 110 ml of TiCl₄ at

120°C for 2 hours. After this period, TiCl₄ was removed by filtration and the solid was then washed with n-heptane at temperatures decreasing from 90°C to room temperature until the chlorine ions disappeared from the filtrate and then kept in heptanic solution.

The catalytic components thus prepared showed the same X-ray spectra similar to those of the catalytic component of Examples 1-10.

The esters, the characteristics of the solid catalytic component and the results of the polymerization tests have been reported in Table III.

Comparative EXAMPLE 4

Example 19 was repeated but using the solid catalytic component prepared according to Example 3 of Italian Pat.

Appl. 26,908 A/78. The X-ray spectrum of that catalytic component was similar to those of Examples 1-10. The characteristics of the solid product and the results of the polymerization test have been reported in Table III.

EXAMPLES 27-29

It was repeated Example 1, but using a solid catalytic component prepared according to Example 1 of German Pat.

Appl. 3,022,738, wherein, instead of ethylbenzoate, esters

of phthalic acid, listed in Table IV, were used and the treatment with TiCl₄ was effected at 120°C.

The X-ray spectrum of the catalytic components was similar to those of Examples 1-10.

The type of ester and the ratios have been reported in Table IV.

The working conditions and the results of the polymerization tests have been reported in Table IV.

Comparative EXAMPLE 5

Example 27 was repeated but using a solid catalytic component prepared according to Example 1 of German Pat. Appl. 3,022,738.

The working conditions and the results of the polymerization test have been reported in Table 3.

EXAMPLES 30 - 34

Example 8 was repeated using instead of phenyltriethoxysilane an equimolecular amount of the alkoxysilanes reported in Table V.

The characteristics of the catalytic components and the results of the polymerization tests have been reported in Table V.

	Solid catalyt	catalytic compone	ent			Polymerization	ation		
Example No.	Ester M	MgCl ₂ ester grinding mols/mols	Ti content of solid component % by weight	Surface area m ² /g	Al(C ₂ H ₅) ₃ PES mols/mols	Catalyst mg	Yield g polymer g catalyst component	H ap	Nin h d1/g
-	d11sobutyl-2,3-naph- thalen-d1carboxylate	14	2.3	ı	10	35	7,750	92.7	1.
2	d1-2-ethyl-hexyl-ma- leate	14	1. 8.	296	10	25	009'9	94	1.1
က	diisobutylmaleate	14	2.9	1	10	54	5,700	92.2	1.2
4	isobutylacetate	7	3.2	í	0.	32	5,700	91.2	1.8
S	isobutylpivalate	7	2.6	1	. 10	22	9,700	92.9	1.3
9	diethil diisobutyl~ malonate		2.4	i	10	47	4,750	94.3	1.7
7	d1-n-butylphthalate	14	2.6	ī	20	35	009'6	91.2	1.3
∞	diisobutylphthalate	14	2.4	350	20	36	16,100	9.1.8	0 <u>0</u> 4
σ	diisobutylphthalate	14	2.4	350	10	38	13,200	95.5	5,9
10	d1-2-ethylhexylsebacate	14	2:3	ŧ	10	52	4,500	94.8	7 <u>7</u>
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TABLE 1

	Solid catalytic component	ytic compon	lent			Polymerization	sation		
Example No.	Ester	MgCl ₂ ester yrinding mols/mols	Ti content of solid component % by weight	i	Surface Al(C2H5)3 area m ² /g PES mols/mols	catalyst 'my	Yield g polymer g catalyst component	1	1.1. h in h 8 d1/g
=	d1-1sobutyl-phthalate	8	3.0	ŧ	10	17	15,000	97.5	1.6
12	d1-isobutyl-phthalate	ھ	1.4	21.8	10	14	7,000	97.2	1.5
comp. 1	ethylbenzoate	7	2.0	266	10	46	000'9	9.06	1.4
comp. 2	ethylbenzoate	7	2,0	266	ĸ	47	4,000	92.7	1.6
comp. 3	ethylbenzoate	7	1.5	1	3,33	09	2,400	91.3	1.2

TABLE II

	Solid catalytic component	tic compor	nent			Polymerization	ation		
Example No.	Ester	MgCl ₂ ester grinding mols/mols	Ti content of solid component g	Surface area m ² /g	Al(C ₂ H ₅) ₃ PES mols/mols	Catalyst	Yield g polymer g catalyst component	I.I.	n in h d1/g
13	diphenylcarbonate	8	2	i	10	29	6,050	93.2	1.5
14	1,2-diidroxy-diacetate benzene	9	1.3	1	10	41	6,100	92.1	1.4
15	ethyl-diphenylacetate	S	1.9	i	10	38	6,400	91.1	1.2
16	diisobutyl-phthalate	7	1.4	157	20	19	9,100	86	1.3
11	diisobutyl-phthalate	7	1.4	157	100	20	000'6	94.3	1.1
18	<pre>1sobuty1-benzoylace- tate</pre>	7	2.1	1	10	42	2,900	95.4	1.2

	Solid catalyt	catalytic component		PoJ	Polymerization		
Example No.	Ester	Ti content of solid component % by weight	Al(C ₂ H ₅) PES mols/mols	Catalyst my.	Yield g polymer g catalyst component	н • œ	11n h d1/9
19	dioctylphthalate	2.7	20	19	12,100	92.2	1.8
20	diisobutylphthalate	2.4	20	22	13,900	6.96	1.2
21	dineopentylphthalate	3.5	20	15	18,500	9.7	1.2
22	diphenylphthalate	2.6	10	25	11,000	94.5	1.8
23	diethyl-butylmalonate	3.9	10	29	12,800	93.3	1.7
24	benzyl butylpthalate	3.1	٤ 50	24	14,000	96.1	1.4
25	mono-2-ethyl-hexylmaleate	2.8	10	20	8,000	92.5	1.3
26	monoisobutylphthalate	3.1	10	24	10,000	94.3	1.6
comp. 4	ethylbenzoate	3.8	ហ	38	6,850	91,5	÷.
							45977 - 2
							9 ° -

TABLE IV

	Solid cata	Solid catalytic component	ent		Polym	Polymerization		
Example No.	Ester	MgCl ₂ ester grinding mols/mols	Ti content of solid component % by weight	Al(C ₂ H ₅) ₃ PES mols/mols	Catalyst	Yield g polymer g catalyst component	I &	71n h d1/g
27	dioctylphthalate	4.7	1.9	20	27	12,000	93.9	1.5
28	lsobutylphthalate	4.7	3.8	20	32	10,000	96.5	1.4
29	neopentylphthalate	9.4	1.9	20	30	12,800	97.8	1.0
ոթ. 5	mp. 5 ethylbenzoate	ĸ	3.1	10	49	7,900	94.2	1.5

TABLE V

	Poly	lymerization	-	-	
Example No.	Alcoxysilane	Catalyst mg	Yield g polymer g catalyst component	I.I.	h in h d1/g
30	ethyltriethoxysilane	33.4	11,000	89.3	1.2
31	chlorophenyldiethoxysilane	44.6	008'6	91.0	1.3
. 32	diphenyldiethoxysilane	25.1	14,300	92.4	1.0
33	phenyldiethoxysilane	31.2	13,200	91.5	1.3
34	diphenyldimethoxysilane	22.2	18,200	93.1	1.5

CLAIMS

- Al-alkyl compound and a silicon compound containing one or more Si-OR, Si-OCOR or Si-NR₂ bonds (R being a hydrocarbyl radical) to form catalysts for the polymerization of alpha-olefins, comprising an anhydrous Mg dihalide in active form, as essential support, and supported on said dihalide a Ti-halide or a Ti-haloalcoholate and an ester selected from the following groups of compounds:
 - 1.1 Mono and polyesters of saturated polycarboxylic acids wherein at least one of the esteric carbo nyl groups is linked to a tertiary or quaternary carbon atom or to a linear or branched chain of at least 4 carbon atoms;
 - 1.2 Mono and polyesters of unsaturated polycarboxy1
 ic acids wherein at least two carboxy groups
 are linked to vicinal double bond-forming carbon
 atoms and in which at least one of the R hydrocarbyl radicals of the -COOR groups is a branched
 saturated or unsaturated radical with 3 to 20
 carbon atoms or is an aryl or arylalkyl radical

with 6 to 20 carbon atoms.

- 1.3 Mono and diesters of aromatic bicarboxylic acids having the COOH groups in ortho position and wherein at least one of the R hydrocarbyl radicals of the COOR groups contains from 3 to 20 carbon atoms;
- 1.4 Mono and polyesters of aromatic hydroxy compounds containing at least two hydroxyl groups in ortho position;
- 1.5 esters of aromatic hydroxy acids wherein at least a hydroxyl group is in ortho position to the carboxy group;
- 1.6 Esters of saturated or unsaturated carboxylic acids wherein at least one of the R, R' hydrocarbyl radicals of the R COOR' groups is a saturated or unsaturated branched radical containing from 3 to 20 carbon atoms or is an arylalkyl radical with 7 to 20 carbon atoms or R is an aryl radical with 3 to 20 carbon atoms and in which when the R' radical is a linear hydrocarbyl radical, it contains from 3 to 20 carbon atoms.

- 1.7 Esters of carbonic acid of formula CO(OR) in which at least one of the R radicals, which can be the same or different, is a hydrocarbyl radical with 3 to 20 carbon atoms.
- A solid component as defined in claim 1) in which the Mg dihalide is Mg dichloride or Mg dibromide, the Ti halide is a Ti tetrahalide and the ester is selected (from the 1.2, 1.3, 1.6 and 177 groups).
- 3) Catalysts for the polymerization of alpha-olefins comprising the product of reaction of the following components:
 - a) an Al-trialkyl or an Al-alkyl compound containing 2 or more Al atoms linked to each other
 through an oxygen or nitrogen atom or through
 SO₄ or SO₃ groups;
 - b) a silicon compound containing one or more Si-OR, Si-OCOR or Si-NR₂ bond, R being a hydrocarbyl radical;
 - c) a solid component as defined in claims 1) to 2).
- 4) Catalysts as defined in claim 3) in which the silicon compound of component b) is a phenyl di- or
 trialkoxylane or an alkyl di- or trialkoxysilane.

• • •

- 5) Catalysts as defined in claim 4) in which the component c) is the solid component as defined in claim
 2).
- OH Process for the polymerization of alpha-olefins

 CH2=CHR in which R is an alkyl radical with 1 to 4

 C, or an aryl radical and mixtures of said olefins

 with ethylene, characterized in that the polymerizat

 ion process is carried out in liquid phase in presence or not of an inert hydrocarbon solvent or in

 gas phase in presence of a catalyst as defined in

 claims 3) to 5).
 - 7) Polymers of the alpha-olefins obtained with the process of claim 6).

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